## Human Group A Streptococci Virulence Genes in Bovine Group C Streptococci

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Phage-encoded virulence genes of group A strepto-cocci were detected in 10 (55.6%) of 18 isolates of group C streptococci that had caused bovine mastitis. Bovine isolates carried other genetic determinants, such as composite transposon Tn1207.3/Φ10394.4 (100%) and antimicrobial drug resistance genes *erm*(B)/*erm*(A) (22.2%), *lin*B (16.6%), and *tet*(M)/*tet*(O) (66.7%), located on mobile elements.

Strains of Streptococcus dysgalactiae subsp. dysgalactiae are described as  $\alpha$ -hemolytic or nonhemolytic (Lancefield group C) and associated only with animal infections (bovine mastitis), a disease with major economic consequences for the dairy industry (1). Group A streptococci (GAS)-specific phage-associated virulence determinants encoding pyrogenic exotoxins or superantigens (speM, ssa), which are strongly associated with severe diseases such as scarlet fever, streptococcal toxic shock syndrome, and rheumatic fever, have been described among human group C streptococci (GCS) or group G streptococci (GGS) (S. dysgalactiae subsp. equisimilis) (2) but not among α-hemolytic GCS (S. dysgalactiae subsp. dysgalactiae) of bovine origin. In contrast, M protein or M-like proteins were found in human GGS/GCS (S. dysgalactiae subsp. equisimilis) and in animal GCS (S. dysgalactiae subsp. dysgalactiae) but only in β-hemolytic strains (3).

Composite transposons and other genetic determinants also considered to be located in specific mobile elements such as macrolide (either encoding methylases [erm genes] or efflux pumps [mef genes]) and tetracycline resistance determinants (tet genes) have been found among streptococcal species of human origin. We studied a collection of field isolates of bovine GCS S. dysgalactiae subsp. dysgalactiae

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to search for genetic determinants, particularly those carried by mobile elements known to be transferred among human GAS and GGS/GCS.

## The Study

We studied 18 α-hemolytic *S. dysgalactiae* subsp. *dysgalactiae* field isolates of Lancefield group C that had caused bovine subclinical mastitis. Isolates were obtained from 304 milk samples of 248 cows from 8 farms in Portugal that were included in the study. Detailed information regarding isolation methods and identification of field isolates by biochemical methods was described in a study of the subclinical mastitis–associated pathogen *S. uberis* (4). To confirm identification of *S. dysgalactiae* subsp. *dysgalactiae*, the 16S rRNA gene was amplified by PCR and sequenced (5). *Smal/cfr*9I-digested DNA banding patterns were obtained by pulsed-field gel electrophoresis for clone identification as described (4).

All genes analyzed by PCR are shown in the online Appendix Table (available from www.cdc.gov/EID/content/16/1/116-appT.htm). The *emm* gene subtyping was performed as described (www.cdc.gov/ncidod/biotech/strep/M-ProteinGene\_typing.htm). Primers used and conditions for PCR were essentially as described elsewhere (online Appendix Table).

Samples without DNA and strains lacking (negative) or carrying (positive) specific genes were used as controls in the PCR. Results were consistent in 2 or 3 PCRs that included these controls. Sequencing of all virulence gene amplicons was performed with the same primers used for amplification (STAB-Vida, Lisbon, Portugal). All sequences were compared with sequences in GenBank by using the BLAST alignment tool (www.ncbi.nlm.nih.gov/BLAST).

Antimicrobial drug resistance against macrolides (erythromycin), lincosamides (pirlimycin), and tetracycline was determined as described (10). Macrolide resistance phenotypes identified were M (resistance to macrolides) and MLS $_{\rm B}$  (resistance to macrolides, lincosamides and streptogramins B).

We detected bacteriophage-associated virulence genes *spe*M, *spe*K, *spe*C, *spd*1, and *spe*L. Overall, *spe*M was found in 10 (55.6%) of 18 bovine GCS isolates, *spe*K in 9 (50%), *spe*C and *spd*1 in 6 (33%), and *spe*L in 4 (22.2%). All but 1 of the PCR products showed expected sizes (online Appendix Table). Tn*1207.3*/Φ10394.4 composite transposon left junction amplicon showed a size of 380 bp instead of 453–6,807 bp as described for GAS (*9*). No amplification was observed for the right junction of this genetic element.

The *emm* gene encoding the antiphagocytic M surface protein was not amplified in any of the 18 bovine GCS isolates; therefore, no *emm* types were obtained. Subsets of isolates were erythromycin and pirlimycin resistant (MLS<sub>R</sub>)

phenotype) and contained *erm*(B) or *erm*(A) genes (22.2%) or erythromycin susceptible and pirlimycin resistant and contained the *linB* gene (16.6%). All isolates were tetracycline resistant with a subset (66.7%) carrying *tet*(M) or *tet*(O) tetracycline resistance determinants. Distribution of bacteriophage-associated virulence genes and other characteristics of strains are shown in Figure 1.

Sequences of all virulence genes were compared by using the BioEdit sequence alignment editor (www. mbio.ncsu.edu/BioEdit/bioedit.html). One different allele was found for each of the following gene sequences: spd1 (among 6 strains), speC (among 6 strains), and speL (among 4 strains). Two alleles were found for speK (among 9 strains) (speK-1 and speK-2), and 4 alleles were found for speM gene sequences (among 10 strains) (speM-1, speM-2, speM-3, and speM-4). Bovine alleles had sizes of 386 bp (spd1), 222 bp (speC), 444 bp (speL), 232 bp (speK), and 357 bp (speM). Examples of alignments between bovine virulence gene alleles with sequences from GenBank (only most similar ones) are shown in Figure 2.

## **Conclusions**

Using PCR, we determined that bovine GCS S. dysgalactiae subsp. dysgalactiae strains (55.6%) carried  $\geq 1$  GAS-specific bacteriophage virulence-associated genes (spd1, speC, speK, speL, and speM). This finding suggested that bacteriophages may also play a role in the genetic plasticity and virulence of animal GCS.

The *spe*L allele from bovine strains showed higher similarity with the *sze*M allele (99% maximum identity) from *S. equi* subsp. *zooepidemicus* than with the *spe*L allele (97% maximum identity) from *S. pyogenes*. The *sze*M gene encodes a superantigen in *S. equi* subsp. *zooepidemicus*, which is primarily a pathogen of nonhuman animal species. This organism causes mastitis in cows and mares and is most frequently found in horses (14). We also observed that 3 of the *spe*M alleles found among bovine strains (*spe*M-1, *spe*M-2, and *spe*M-3) also showed higher similarity with superantigen-encoding gene *sze*L from *S. equi* subsp. *zooepidemicus* than with *spe*M gene sequence from *S. pyogenes*. Another allele (*spe*M-4) showed higher similarity with the *sdm* gene from *S. dysgalactiae* subsp. *dysgalactiae* than with the speM gene from *S. pyogenes*.

The remaining alleles (spd1, speC, speK-1, and speK-2) from the GCS S. dysgalactiae subsp. dysgalactiae bovine strains showed high similarity with S. pyogenes superantigen genes (98%–99% maximum identity). This finding supports our hypothesis that GAS prophages may play a role in the genetic plasticity of this pathogen. The speC and spd1 genes are known to be localized on the same GAS prophage (15), and both genes were detected in 6 bovine GCS S. dysgalactiae subsp. dysgalactiae isolates in our study.

None of 18 α-hemolytic group C S. dysglacatiae subsp. dysgalactiae bovine isolates in this study were typed by emm-typing because amplification products in the PCR

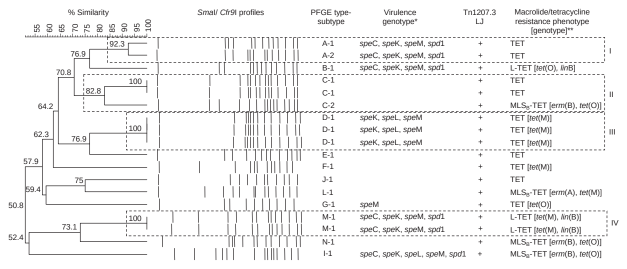


Figure 1. Dendrogram and pulsed-field gel electrophoresis (PFGE) profiles of group C streptococci (*Streptococcus dysgalactiae* subsp. *dysgalactiae*) subclinical mastitis isolates from 8 dairy herds, Portugal. PFGE type-subtype, virulence genotype, antimicrobial drug resistance phenotypes, and genotypes of each isolate are indicated. The dendrogram was produced by using Dice coefficients and unweighted pair group method using arithmetic averages. Default clustering settings of 0.00% optimization (i.e., the relative distance an entire lane is allowed to shift in matching attempts) and 1.5% band position tolerance were used. \*All isolates were negative for *speA*, *ssa*, *speH*, *speJ*, *speI*, and *slaA* genes and for Tn 1207.3/Φ10394.4 element right junction tested by PCR; \*\*All isolates were negative for *mefA*, *tet*(T), *tet*(U), *tet*(Q), *tet*(S) and *tet*(K) genes tested by PCR; TET, resistance only to tetracycline; MLS<sub>B</sub>-TET, resistance to macrolides, lincosamides, streptogramin B, and TET; L-TET, susceptibility to macrolides and resistance to lincosamides (L phenotype) and TET; Tn1207.3 LJ, Tn1207.3/Φ10394.4 element left junction. Clusters are shown in roman numerals on the right.

spd1-bovine spd1-MGAS9429	345 ATTGTCACTA ATTGCCACTA	455 ATAGTT <b>A</b> TGC ATAGTT <b>G</b> TGC	515 TTATCTTGC TCTATCTTGC			
speK-1-bovine speK-2-bovine speK-MGAS315	75 TCTTTTCAGA TCTTTTTCAGA TCTTTTCAGA	135 GATATGAT <b>G</b> A GATATGAT <b>A</b> A GATATGAT <b>A</b> A				
speC-bovine speC-MGAS9429	545 GTAAG <b>T</b> GGCA GTAAG <b>C</b> GGCA					
speL-bovine szeM-S.equi.zoo.	255 TTATTAT <b>G</b> GA TTATTAT <b>A</b> GA	365 AATGTCGAT <b>A</b> AATGTCGAT <b>G</b>	595 ATTTATGC <b>A</b> G ATTTATGC <b>T</b> G		A	
speL-bovine speL-MGAS8232	325 CCTATACCCA CCTACACCCA	365 AATGTCGATA AATGTCGATG	385 TGCATTATCG TGCATTATCA	465 GAGCTGGAAA GAGCCGGAAA	505 GCATAAGAAA GCATAAAAAA	515 GATGAGACCT GATGATACCT
speL-bovine speL-MGAS8232	565 CCAGGA <b>G</b> GTT CCAGGA <b>A</b> GTT	595 ATTTATGCAG ATTTATGCTG	625 TCAGTATGGT TCAGTATGAT			
speM-1-bovine speM-2-bovine speM-3-bovine szeL-S.equi.zoo.	345 AGGAGGAGA AGGAGGAGA AGGAGGAGGA TTGGTGATCA	355 GGTTGATATT GGTTGATATT GGTTGATATT T GTGAATATT	365 TATGCTCTAG TATGCTCTAG TATGCTCTAG TATGCTCTAG	385 TGATGTTGTA TGATGTTGTA TGATGTTGTA TCATGTTGTA	575 AGGAAAAGTT AGGAAAAGTA AGGAAAAGTT AGGAAAAGTT	665 GGTGGCACTA GGTGGCACTA GGTGGCATTA GGTGGCATTA
SpeM-4-bovine speM-MGAS8232	275 AGGA <b>G</b> G TTTAAGGA <b>A</b> G	285 A <b>G</b> GA <b>G</b> GTTGA A <b>A</b> GA <b>A</b> GTTGA	365 ATAA <b>A</b> AACAA ATAA <b>G</b> AACAA	385 AAAACTTGAT GAAACTTGAT	585 TTCATTACAA TTCATTATAA	625 CACCCATCCT CACTCATCCT

Figure 2. Alignments of bovine group C streptococci (Streptococcus dysgalactiae subsp. dysgalactiae) alleles of virulence genes from 8 dairy herds, Portugal, with sequences from the National Center for Biotechnology (Bethesda, MD, USA) database showing base differences between sequences. The alignments were created by using BioEdit sequence alignment editor (www.mbio.ncsu. edu/BioEdit/bioedit.html). Nucleotide differences are shown in **boldface**. A) spd1 (99% maximum identity); B) speK (99% maximum identity); C) speC (99% maximum identity); D) speL-szeM (99% maximum identity); E) speL (97% maximum identity; F) speM alleles 1, 2, and 3-szeL (98%-99% maximum identity); G) speM allele 4 (98% maximum identity). S. equi. zoo., S. equi subsp. zooepidemicus.

specific for the M surface protein gene *emm* were not obtained. This result is consistent with those of a report that  $\beta$ -hemolytic, but not  $\alpha$ -hemolytic, group C *S. dysglacatiae* subsp. *dysgalactiae* isolates of animal origin contained the *emm* gene (3).

Amplification (380-bp product) of the left junction of the composite transposon in bovine isolates suggests that this mobile element may be inserted in a similar location, the *comEC* locus, as mapped in *S. pyogenes* and *S. dysglactiae* subsp. *equisimilis*. Absence or unexpected PCR products specific for any of the junctions of this element have been reported in other studies and attributed to possible lack of homology between the target and primers used (9). Detection of the *linB* gene carried by a large conjugative plasmid (13) in 3 of 18 bovine GCS *S. dysgalactiae* subsp. *dysgalactiae* isolates is indicative of horizontal gene transfer.

Our findings indicate that  $\alpha$ -hemolytic bovine GCS isolates, which are known to be environmental or contagious pathogens and a cause of bovine mastitis, may be reservoirs of virulence genes encoded by prophages of human-specific GAS. These genes encode exotoxins, superantigens, and streptodornases, which are responsible for GAS virulence and pathogenesis, and may be transferred to other streptococci of human origin by horizontal genetic transfer. Therefore,  $\alpha$ -hemolytic isolates should not be disregarded as putative infectious disease agents in humans.

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Ms Rato is a doctoral candidate in the Department of Life Sciences of the Faculty of Science and Technology, New University of Lisbon, Caparica, Portugal. Her research interests include epidemiology, antimicrobial drug resistance, and virulence mechanisms of streptococci from animal and human origin.

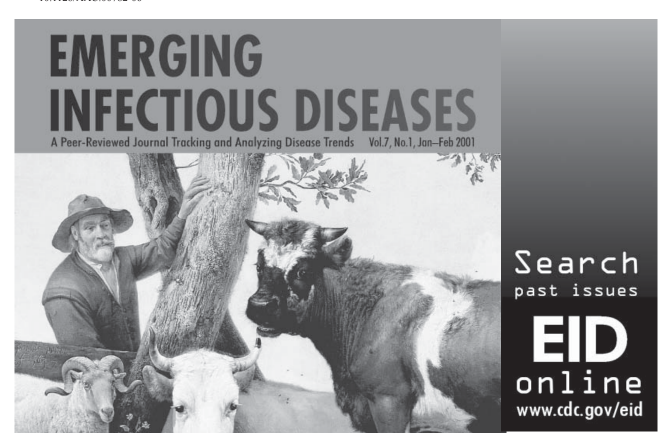
## References

- Vieira VV, Teixeira LM, Zahner V, Momen H, Facklam RR, Steigerwalt AG, et al. Genetic relationships among the different phenotypes of *Streptococcus dysgalactiae* strains. Int J Syst Bacteriol. 1998;48:1231–43.
- Igwe EI, Shewmaker PL, Facklam RR, Farley MM, van Beneden C, Beall B. Identification of superantigen genes speM, ssa, and smeZ in invasive strains of beta-hemolytic group C and G streptococci recovered from humans. FEMS Microbiol Lett. 2003;229:259–64. DOI: 10.1016/S0378-1097(03)00842-5

- Zhao J, Hayashi T, Saarinen S, Papageorgiou AC, Kato H, Imanishi K, et al. Cloning, expression, and characterization of the superantigen streptococcal pyrogenic exotoxin G from *Streptococcus dysgalactiae*. Infect Immun. 2007;75:1721–9. DOI: 10.1128/IAI.01183-06
- Rato MG, Bexiga R, Nunes SF, Cavaco LM, Vilela CL, Santos-Sanches I. Molecular epidemiology and population structure of bovine *Streptococcus uberis*. J Dairy Sci. 2008;91:4542–51. DOI: 10.3168/jds.2007-0907
- Weisburg WG, Barns SM, Pelletier DA, Lane DJ. 16S ribosomal DNA amplification for phylogenetic study. J Bacteriol. 1991;173:697–703.
- Pires R, Rolo D, Mato R, Feio de Almeida J, Johansson C, Henriques-Normark B, et al. Resistance to bacitracin in *Streptococcus pyogenes* from oropharyngeal colonization and noninvasive infections in Portugal was caused by two clones of distinct virulence genotypes. FEMS Microbiol Lett. 2009;296:235–40. DOI: 10.1111/j.1574-6968.2009.01642.x
- Lintges M, Arlt S, Uciechowski P, Plümäkers B, Reinert RR, Al-Lahham A, et al. A new closed-tube multiplex real-time PCR to detect eleven superantigens of *Streptococcus pyogenes* identifies a strain without superantigen activity. Int J Med Microbiol. 2007;297:471–8. DOI: 10.1016/j.ijmm.2007.03.015
- Green NM, Beres SB, Graviss EA, Allison JE, McGeer AJ, Vuopio-Varkila J, et al. Genetic diversity among type emm28 group A *Streptococcus* strains causing invasive infections and pharyngitis. J Clin Microbiol. 2005;43:4083–91. DOI: 10.1128/JCM.43.8.4083-4091.2005
- Figueiredo TA, Aguiar SI, Melo-Cristino J, Ramirez M. DNA methylase activity as a marker for the presence of a family of phage-like elements conferring efflux-mediated macrolide resistance in streptococci. Antimicrob Agents Chemother. 2006;50:3689–94. DOI: 10.1128/AAC.00782-06

- Pires R, Rolo D, Gama-Norton L, Morais A, Lito L, Salgado MJ, et al. Group A streptococci from carriage and disease in Portugal: evolution of antimicrobial resistance and T antigenic types during 2000–2002. Microb Drug Resist. 2005;11:360–70. DOI: 10.1089/ mdr.2005.11.360
- Ng L-K, Martin I, Alfa M, Mulvey M. Multiplex PCR for the detection of tetracycline resistant genes. Mol Cell Probes. 2001;15:209–15. DOI: 10.1006/mcpr.2001.0363
- Aminov RI, Garrigues-Jeanjean N, Mackie RI. Molecular ecology of tetracycline resistance: development and validation of primers for detection of tetracycline resistance genes encoding ribosomal protection proteins. Appl Environ Microbiol. 2001;67:22–3. DOI: 10.1128/AEM.67.1.22-32.2001
- Bozdogan B, Berrezouga L, Kuo MS, Yurek DA, Farley KA, Stockman BJ, et al. A new resistance gene, *linB*, conferring resistance to lincosamides by nucleotidylation in *Enterococcus faecium* HM1025. Antimicrob Agents Chemother. 1999;43:925–9.
- Alber J, El-Sayed A, Estoepangestie S, Lämmler C, Zschöck M. Dissemination of the superantigen encoding genes seeL, seeM, szeL and szeM in Streptococcus equi subsp. equi and Streptococcus equi subsp. zooepidemicus. Vet Microbiol. 2005;109:135–41. DOI: 10.1016/j.vetmic.2005.05.001
- Banks DJ, Beres SB, Musser JM. The fundamental contribution of phages to GAS evolution, genome diversification and strain emergence. Trends Microbiol. 2002;10:515–21. DOI: 10.1016/S0966-842X(02)02461-7

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Appendix Table. Genes analyzed by PCR and primers used, for group C streptococci, from 8 dairy herds, Portugal

Primer	Sequence (5´? 3´)	Expected product, bp	Reference
Prophage-associated virulence determinants			
Pyrogenic exotoxins			
ssa (forward)	GTGTAGAATTGAGGTAATTG	706	( <u>6</u> )
ssa (reverse)	TAATATAGCCTGTCTCGTAC		
speA (forward)	CTTAAGAACCAAGAGATGGC	200	( <u>6</u> )
speA (reverse)	ATAGGCTTTGGATACCATCG		
speC (forward)	CATCTATGGAGGAATTACGC	246	( <u>6</u> )
speC (reverse)	TGTGCCAATTTCGATTCTGC		
speH (forward)	AGATTGGATATCACAGG	416	( <u>6</u> )
speH (reverse)	CTATTCTCTCGTTATTGG		
spel (forward)	AAGGAAAATAAATGAAGGTCCGCCAT	217	( <u>Z</u> )
spel (reverse)	TCGCTTAAAGTAATACCTCCATATGAATTCTTT		
speJ (forward)	ATCTTTCATGGGTACG	535	( <u>6</u> )
speJ (reverse)	TTTCATGTTTATTGCC		
speK (forward)	TATCGCTTGCTCTATACACTACTGAGAGT	233	( <u>Z</u> )
speK (reverse)	CCAAACTGTAGTATTTTCATCCGTATTAAA		
speL (forward)	GGACGCAAGTTATTATGGATGCTCA	460	( <u>7</u> )
speL (reverse)	TTAAATAAGTCAGCACCTTCCTCTTTCTC		
speM (forward)	GCTTTAAGGAGGAGGAGGTTGATATTTATGCTCTA	411	( <u>7</u> )
speM (reverse)	CAAAGTGACTTACTTTACTCATATCAATCGTTTC		
DNase 1			
spd1 (forward)	CCCTTCAGGATTGCTGTCAT	400	( <u>8</u> )
spd1 (reverse)	ACTGTTGACGCAGCTAGGG		
Phospholipase A2e			
slaA (forward)	CTCTAATAGCATCGGCTACGC	440	( <u>8</u> )
slaA (reverse)	AATGGAAAATGGCACTGAAAG		
Composite transposon			
Tn 1207.3/F10394.4 RJ* (forward)	CGAGGAGTTAGTATGGAAAC	473	( <u>9</u> )
Tn 1207.3/F10394.4 RJ* (reverse)	CCCATAATAGGCAACTGGTCTCCAGC		
Tn 1207.3/F10394.4 LJ* (forward)	TCTTCGCCGCATAAACCCTATC	453/6,807†	( <u>9</u> )

CCTTTGACCAATGAAGTGACCTTT

Tn 1207.3/F10394.4 LJ\* (reverse)

Antimicrobial drug resistance determinants

Macrolide resistance			
mef(A) (forward)	GACCAAAAGCCACAATTGTGGA	1,432	( <u>10</u> )
mef(A) (reverse)	CCTCCTGTCTATAATCGCATG	., .==	( <u></u> )
erm(A) [subclass erm(TR)] (forward)	CCCGAAAAATACGCAAAATTTCAT	590	( <u>10</u> )
erm(A) [subclass erm(TR)] (reverse)	CCCTGTTTACCCATTTATAAACG		(==)
erm(B) (forward)	GGAGTGATACATGAACAAAATA	531	( <u>10</u> )
erm(B) (reverse)	TTCCTTTTAGTAACGTGTAACTTT		(/
Tetracycline resistance			
tet(M) (forward)	TGGAATTGATTTATCAACGG	1,080	( <u>10</u> )
tet(M) (reverse)	TTCCAACCATACAATCCTTG		` ,
tet(O) (forward)	AACTTAGGCATTCTGGCTCAC	515	( <u>11</u> )
tet(O) (reverse)	TCCCACTGTTCCATATCGTCA		
tet(T) (forward)	AAGGTTTATTATAAAAGTG	169	( <u>12</u> )
tet(T) (reverse)	AGGTGTATCTATGATATTTAC		
tet(W) (forward)	GAGAGCCTGCTATATGCCAGC	168	( <u>12</u> )
tet(W) (reverse)	GGGCGTATCCACAATGTTAAC		
tet(Q) (forward)	TTATACTTCCTCCGGCATCG	904	( <u>11</u> )
tet(Q) (reverse)	ATCGGTTCGAGAATGTCCAC		
tet(S) (forward)	GAAAGCTTACTATACAGTAGC	169	( <u>12</u> )
tet(S) (reverse)	AGGAGTATCTACAATATTTAC		
tet(L) (forward)	TCGTTAGCGTGCTGTCATTC	267	( <u>11</u> )
tet(L) (reverse)	GTATCCCACCAATGTAGCCG		
tet(K) (forward)	TCGATAGGAACAGCAGTA	169	( <u>11</u> )
tet(K) (reverse)	CAGCAGATCCTACTCCTT		
Lincosamide resistance			
lin(B) (forward)	CCTACCTATTGTTTGTGGAA	925	( <u>13</u> )
lin(B) (reverse)	ATAACGTTACTCCCTATTC		
M protein			
emm (forward)	TATT(C/G)GCTTAGAAAATTAA	Variable	‡
emm (reverse)	GCAAGTTCTTCAGCTTGTTT		

<sup>\*</sup>RJ, right junction; LJ, left junction.

<sup>†</sup>The expected ampicon size was 454 bp according to the reported organization of the Tn1207.3 element or 6,807 bp according to the reported sequence of F10394.4 (9).

Human GAS Virulence Genes in Bovine GCS, Appendix Table   CDC EID				
‡www.cdc.gov/ncidod/biotech/strep/M-ProteinGene typing.htm				